

# WORLD INTELLECTUAL PROPERTY ORGANIZATION



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(54) Tide: HETEROVESICULAR LIPOSOMES



### (57) Abstract

Disclored are heterovolander liprocomes containing different biological compositions (14, 14a) such enapsulated in reparate biometers of the liprocomes, having defined size distribution, adjustable internal chamber size and rounder, method of making them and treatment of planties with them. The preparation process behavior het defined of composition for limit just component in the vidal to obtain an emislate, the additions of composition to ascend lipid component in the vidal to obtain a second emislate and rising between emislates of them as obtained.

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### HETEROVESICULAR LIPOSOMES

# Field of the Invention

The invention relates to the synthetic heterovesicular lipid vesicles or liposomes, processes for their manufacture and encapsulation of various materials therein, and treatment of patients with them.

# Background Art Multivesicular liposomes are one of the three main types of liposomes, first made by Kim, et al. (1983,

Biochim, Biophys. Acta 782, 339-348), and are uniquely different from the unilamellar (Huang, 1969, Biochemistry 8,344-352; Kim, et al. 1981, Biochim. Biophys. Acta 646, 1-10) and multilamellar (Bangham, et al. 1965, J. Mol. Bio. 15 13,238-252) liposomes in that there are multiple nonconcentric aqueous chambers within. Previously described techniques for producing liposomes relate to the production of non-multivesicular liposomes; for example, U.S. Patent Nos. 4,522,803 - Lenk, 4,310,506 - Baldeschwieler,

4,235,871 - Papahadjopoulos, 4,224,179 - 4,078,052 -Papahadjopoulos, 4,394,372 - Taylor, 4,308,166 - Marchetti, 4,485,054 - Mezei, and 4,508,703 - Redziniak. For a comprehensive review of various methods of liposome preparation, refer to Szoka, et al. 1980, Ann. Rev. Biophys. 25 Bioeng. 9:467-508.

Heterovesicular liposomes are lipid vesicles or liposomes with multiple internal aqueous chambers where at least two substances of different compositions are each encapsulated in separate chambers within one liposomes. The lipid vesicles or liposomes with multiple internal aqueous

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chambers include, but are not limited to, multilamellar liposomes, stable pausilamellar liposomes, under multivesicular liposomes. It is highly advantageous to provide a liposome delivery system in which two or more different substances are each encapsulated in separate compartments of a single liposome rather than encapsulated towether in each compartment of the liposome.

## Summary of the Invention

The composition of the present invention comprises heterovesicular liposomes, i.e. lipid vesicles or liposomes with multiple internal aqueous chashers where two or more substances of different compositions are each encapsulated separately in different chambers within one liposome.

Briefly, the method of the invention comprises

making a "water-in-lipid" emulsion by dissolving amphipathic lipids in one or more organic solvents for the first lipid component, adding an immiscible first aqueous component including a substance to be encapsulated, preferably in the presence of hydrochloric acid, and then emulsifying the mixture mechanically. In the emulsion, the water droplets suspended in the organic solvent will form the internal aqueous chambers, and the monolayer of amphipathic lipids lining the aqueous chambers will become one leaflet of the bilayer membrane in the final product. A second lipid component is then formed by dissolving amphipathic lipids in a volatile organic solvent and adding an immiscible second aqueous component including a second substance to be encapsulated, preferably in the presence of hydrochloric acid. A second emulsion is then created. A chimeric emulsion is then formed by combining the first and second emulsions. The chimeric emulsion consists of multiple water droplets suspended in organic solvent where the substances of two different compositions are each dissolved separately in different aqueous droplets. The chimeric emulsion is then immersed in a third aqueous immiscible component preferably containing one or more nonionic osmotic agents

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and acid-neutralizing agent of low ionic strength and then mechanically dividing it to form solvent spherules suspended in the third aqueous component. The solvent spherules contain multiple aqueous droplets where the substances of two different compositions are each dissolved separately in different aqueous droplets within a single solvent spherule. The volatile organic solvent is evaporated from the spherules preferably by passing a stream of gas over the suspension. When the solvent is completely evaporated, the spherules convert into heterovesicular liposoms with multiple internal aqueous chambers where two substances of different compositions are encapsulated separately in different chambers within one liposom

The use of hydrochloric acid with a neutralizing agent, or other hydrochlorides which slow leakage rates is preferably for high encapsulation efficiency and for a slow leakage rate of encapsulated solicules in biological fluids and in yim. It is also preferable to use neutralizing agent of low ionic strength to prevent solvent spherules from sticking to each other.

Accordingly, it is an object of the present invention to provide a heterovesicular lipid vesicle or liposome having at least two substances of different compositions each encapsulated separately in different chambers of the vesicle or liposomes.

A further object of the present invention is the provision of a heterovesicular liposeme containing at least two biologically active substances of different compositions each encapsulated separately in chambers of the liposome in the presence of hydrochloric acid or other hydrochlorides which slow the leakage of them.

It is a further object of the present invention to provide a heterovesicular liposome containing at least two biologically active substances of different compositions each encapsulated separately in chambers of the liposome in

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the presence of hydrochloric acid or other hydrochlorides and a neutralizing agent.

It is a further object of the present invention to provide methods of producing such heterovesicular lipid vesicles or liposomes.

It is a further object of the present invention to

provide processes for producing such heterovesicular lipid vesicles or liposomes by providing a first lipid component dissolved in one or more organic solvents and adding to the lipid component an immiscible first aqueous component containing a first substance to be encapsulated, forming a first water in oil emulsion from the first two immiscible components, providing a second lipid component dissolved in one or more organic solvents and adding into the lipid component an immiscible second aqueous component containing a second substance to be encapsulated, forming a second water in oil emulsion from the second two immiscible components, forming a chimeric emulsion by combining the first water in oil emulsion and second water in oil emulsion, transferring and immersing the chimeric emulsion into a third immiscible aqueous component, dispersing the chimeric emulsion to form solvent spherules containing multiple droplets of the first aqueous component containing the first substance and the second aqueous component containing the second substance, and evaporating the organic solvent from the solvent spherules to form the heterovesicular lipid vesicles or liposomes.

It is a further object to provide such a process in which a variety of hydrophilic biologically active materials and can be encapsulated separately in chambers of the heterovesicular lipid vesicles or liposomes.

It is a further object of the present invention to provide a method for the treatment of a patient with at least two separate biologically active substances of different compositions by administering them to the patient

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encapsulated separately in chambers of a heterovesicular vesicle or liposome.

Other and further objects, features and advantages of the invention appear throughout the specification and claims.

## Brief Description of the Drawings

Figures 1-8 are schematic diagrams illustrating preparation of a heterovesicular vesicle or liposome.

Description of Preferred Embodiments
The term "multivesicular liposomes" as used

The term "multivesicular liposomes" as used throughout the specification and claims means man-made, microscopic lipid-vesicles consisting of lipid bilayer membranes, enclosing multiple non-concentric aquecus chambers which all contain the same component. In contrast, the term "heterowssicular liposomes as used throughout the specification and claims means man-made, microscopic liquid vesicles consisting of lipid bilayer membranes enclosing multiple, aquecus chamber wherein at least two of the chambers separately contain substances of different compositions. The microscopic lipid vesicles include but are not limited to multilemellar liposomes, stable puncilemellar liposomes, and multivesicular liposomes.

The term "chimeric emulsion" as used throughout the specification and claims means an emulsion that consists of multiple water droplats suspended in organic solvent where the substances of two different compositions are each dissolved separately in different squeeze droplate.

The term "solvent spherule" as used throughout the specification and claims means a microscopic spheroid droplet of organic solvent, within which is multiple smaller droplets of aqueous solution. The solvent spherules are suspended and totally immersed in a second equeous solution. The term "neutral lipid" means oil or fat that

have no membrane-forming capability by themselves and lack a hydrophilic "head" group.

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The term amphipathic lipids means those molecules that have a hydrophilic "head" group and hydrophobic "tail" group and have membrane-forming capability.

The composition of the present invention is a heteroveaicular lipid vesicle or liposome having at least two substances of different compositions each encognulated separately in different chambers of the vesicle or liposome. Many and varied biological substances can be

incorporated by encapsulation within the multivesicular liposomes. These include drugs, and other kinds of materials, such as DNA, BNA, proteins of various types, protein hormones produced by recombinant DNA technology effective in humans, hematopoietic growth factors, monokines, lymphokines, tumor necrosis factor, inhibin, tumor growth factor alpha and beta, mullerian inhibitory substance, nerve growth factor, fibrohlast growth factor, platelet-derived growth factor, pituitary and hypophyseal hormones including LH and other releasing hormones, calcitonin, proteins that serve as immunosem for

vaccination, and DNA and RNA sequences.

The following Table 1 includes a list of representative biologically active substances which can be encapsulated in heterovesicular liposomes in the presence of a hydrochloride and which are effective in humans.

25	TABLE 1			
	Antiasthma	Antiarrhythmic	Tranquilizers	
	metaproterenol	propanolol	chlorpromazine	
	aminophylline	atenolol	benzodiazepine	
	theophylline	verapamil	butyrophenones	
30	terbutaline	captopril	hydroxyzines	
	Tegretol	isosorbide	meprobamate	
	ephedrine		phenothiazines	
	isoproterenol		reservine	
	adrenalin		thioxanthines	
35	norepinephrine			

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	Cardiac glycosides	Hormones	Steroids
	digitalis	antidiuretic	prednisone
	digitoxin	corticosteroids	triamcinolone
	lanatoside C	testosterone	hydrocertisone
5	digoxin	estrogen	dexamethasone
		thyroid	betamethosone
		growth	prednisolone
		ACTH	preditacione
		progesterone	
10		gonadotropin	
		mineralocorticoid	
		LH	
		LHRR	
		PSH	
15		calcitonin	
	Antihypertensives	Antidiabetic	Antihistamines
	apresoline	Diabenese	pyribenzamine
	atenolol	insulin	chlorphoniramine
			diphenhydramine
			- Promiyor antitio
20	Antiparasitic	Anticancer	Sedatives & Analgesic
	praziquantel	azathioprine	morphine
	metronidazole	bleomycin	dilandid
	pentamidine	cyclophosphamide	codeine
		adriamycin	codeine-like synthetics
25		daunorubicin	damerol
		vincristine	oxymorphone
		methotrexate	phenobarbital
		6-TG	barbiturates
		6-MP	
30		vinblastine	
		VP-16	
	100	V16-26	
		cisplatine	
		FU	
35	Antibiotic	Immunoptherapies	Vaccines
	penicillin	interferon	influenza
	tetracycline	interleukin-2	respiratory syncytial
	erythromycin	monoclonal antibodies	virus
	cephalothin	camerlobulin	Hemorahi luga indlusees
40		gazmaglobulin	Hemophilus influenza
40	cephalothin	gazmaglobulin	Hemophilus influenza vaccine

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Antibiotic (continued) Antifungal vancomycin amphotericin B gentamycin myconazole muramyl dipeptide tobramycin piperacillin clotrimazole moxalactam amovicillin ampicillin Antihypotension cefazolin dopamine 10 cefadroxil dextroamphetamine cefoxitin other aminoglycosides Proteins and Glycoproteins lymphokines 15 interleuking - 1, 2, 3, 4, 5, and 6 cvtokines GM-CSF M-CSF G-CSF 20 tumor necrosis factor inhihin tumor growth factor Mullerian inhibitors substance nerve growth factor 25 fibroblast growth factor platelet derived growth factor coagulation factors (e.g. VIII, IX, VII) insulin tissue plasminogen activator 30 histocompatibility antigen oncogene products myelin basic protein collagen fibronectin 35 laminin

other proteins made by recombinant DNA

technology

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acyclovir and derivatives Winthroo-51711

Antiviral

ribavirin rimantadine/amantadine azidothymidine & derivatives

adenine arabinoside amidine-type protease inhibitors

Other cell surface receptor

Nucleic Acids & Analogs DNA RNA methylphosphonates and analogs

A preferred method of making the heterovericular vesicle or liposome is illustrated in the draving to which reference is now made. In step 1 (Figure 1) a first aqueous substance of composition 10 to be encepsulated is added to a

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first lipid component 12 in the vial 14. The vial 14 is sealed and in step 2 (Figure 2) is mixed and shaken, such as being attached to the head of a vortex mixer to form the first water in oil emulsion 16 containing the first substance of composition 10 to be encapsulated. In step 3

first water in oil emulsion 16 containing the first substance of composition 10 to be encapsulated. In step 3 (Figure 3), a second vial 14s, a second aqueous 10s to be encapsulated is added to a second lipid component 12s, and the vial 14s is sealed and in step 4 (Figure 4) is mixed, such as being attached to the head of a vortex mixer to form a second water-in-oil emulsion 16s containing the substance of composition 10s to be encapsulated.

In step 5 (Figure 5) the first 16 and second 16a water in oil emulsions are added together and mixed, such as by hand to make a "chimeric" emulsion.

In step 6 (Figure 6) a portion of the chimeric enulsion from step 5 is individually added to vials containing a third immiscible aqueous component 18s such as by squirting rapidly through a narrow tip pasteur pipette into two one-dram vials. here shown as one.

20 In step 7 (Figure 7) vials from step 6 are shaken, such as by a vortex mixer, and in step 8 (Figure 8) the chloroform spherule suspension in each vial is transferred from step 7 and the chloroform is evaporated, such as by a stream of nitrogen gas, thereby providing the

25 heterovesicular liposome that contains a first substance in one or more internal aqueous chambers and a second substance in the remaining internal aqueous chambers within a single liposome.

Preferably, each of the substances to be encapsulated are encapsulated in the presence of a hydrochloride, such as hydrochloric acid, which slows their leakage rate from the liposome or vesicle.

As previously mentioned, any biologically active substance, such as illustrated in Table 1, can be encapsulated separately in chambers of the vesicle or licosome.

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The following examples set forth presently preferred methods of encapsulating two substances of different compositions in separate chambers of a vesicle or liposome.

## Example 1

# Preparation of Dideoxycytidine/Glucose Heterovesicular Liposomes

Sign\_1: A first aqueous substance (one ml of 20 mg/ml dideoxycytidine solution in water with 0.1 N hydrochloric acid) was added into a one-dram vial containing the first lipid component (9.3 meales of diolecyl lecithin, 2.1 uncles of diplamitorly phosphatidylglycerol, 15 uncles of cholesterol, 1.8 uncles of triolein and one ml of choleroform).

Step 2: The first vial was sealed and attached to the head of a vortex mixer and shaken at maximum speed for 6 minutes to form the first water-in-oil emulsion.

<u>Step 2</u>: In second vial, the second aqueous substance (one ml of 30 mg/ml glucose solution in water with 0.1 N hydrochloric acid) was added into the second lipid component (which is identical to the first lipid component).

Step 4: The second vial was sealed and attached to the head of a vortex mixer and shaken at maximum speed for 6 minutes to form the second water-in-oil emulsion.

Step 5: 0.5 ml of the first emulsion was added to the second vial and mixed by hand to make a "chimeric" emulsion.

Step 6: Half of the "chimeric" emulsion was individually squirted repidly through a narrow tip Pasteur pipette into one-dram vials, each containing a third immiscible aqueous component (2.5 ml water, 32 mg/ml glucose, 40 mM free-base lysine.

Step\_: The vials from step 6 were shaken on the vortex mixer for 3 seconds at "s" setting to form solvent spherules containing multiple droplets of the first and second aqueous substances within.

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Step 8: The chloroform spherule suspensions in each vials were transferred into the bottom of a 2 L beaker containing 4.5 ml of water, 35 mg/lml glucose, and 22 ml free-base lysine. A stream of nitrogen gas at 7 L/min was flushed through the beaker to evaporate chloroform over 5 minutes at 15 dec. C.

The above example describes a method of making heterovesicular liposomes which separately contain glucose in approximately 5/6 of the internal aqueous chambers and separately contain dideoxycytidine in the remaining 1/6 of the internal aqueous chambers within a single liposome. Heterovesicular liposomes containing dideoxycytidine solution as one aqueous substance and glucose as the second aqueous substance were markedly more stable than non-heterovesicular liposomes.

Example 2

This example is for the synthesis of heterovesicular liposomes containing IL-2 (interleukin-2) and lysine hydrochloride: For each batch of liposomes prepared, one ml of water containing 10 mg/ml HSA (Human serum albumin), 1 ug of IL-2, 200 mM lysine HCl pH 7.13 was added into a one-dram vial containing 9.3 umoles of diolecyl lecithin, 2.1 umoles of dipalmitoyl phosphatidylglycerol, 15 umoles of cholesterol, and 1.8 umoles of triolein and one ml of chloroform (this is the first water-in-oil emulsion). For the second water-in-oil emulsion, 1 ml of lysine HC1 (without IL-2) was added into one-dram vial containing 9.3 umoles of dioleoyl lecithin, 2.1 umoles of dipalmitoyl phosphatidylglycerol, 15 umoles of cholesterol, and 1.87 umoles of triolein and one ml of chloroform. Each of the two vials were individually attached to the head of a vortex mixer and shaken sequentially at the maximum speed for 6 minutes.

0.5 ml of the first water-in-oil emulsion was added to the 2 ml of the second emulsion and mixed to make a "chimeric" water-in-oil emulsion. Half of the "chimeric"

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emilsion was individually squirted rapidly through a narrow tip Pasteur pipetts into one-drraw vials, each containing 2.5 al of 48 glucose in water and 0.1 ml of 1 yains free base, 200 MK, and shaken at maximus speed for 3 seconds to form chloroform spherules. The chloroform spherule suspensions were transferred into 250 ml Erlemmeyer flask containing 5 ml of 44 glucose in water and 0.2 ml of lysine free base, 200 MK. A stream of nitrogen gas at 7 Lymin was flushed through the flask to evaporate chloroform over 5 minutes at 37 decrees C.

### Example 3

This example is for the synthesis of heterovesicular liposomes containing ara-C solution as the first aqueous substance and distilled water as the second aqueous substance. For each batch of liposomes prepared, one ml of water containing 100 mg/ml ara-C, pH 1.1 was added into a one-dram vial containing 9.3 umoles of diolecyl lecithin, 2.1 umoles of dipalmitoyl phosphatidylglycerol, 15 umoles of cholesterol, and 1.8 umoles of triolein and one ml of chloroform, attached to the head of the vortex mixer and shaken at maximum speed for 6 minutes (this is the first water-in-oil emulsion). For the in situ generation of the second water-in-oil emulsion, 1/2 of the content was removed from the first water-in-oil emulsion, and then 1 ml of distilled water was added into the remaining first water-inoil emulsion and the one-dram vial was shaken for 10 seconds at maximum speed. This resulted in a "chimeric" water-inoil emulsion. Half of the "chimeric" emulsion was individually squired rapidly through a narrow tip Pasteur pipette into one-dram vials, each containing 2.0 ml of 4% glucose in water and 0.5 ml of lysine free base, 200 mM, and shaken at maximum speed for 3 seconds to form chloroform spherules. The chloroform spherule suspensions were transferred into 250 ml Erlenmeyer flask containing 4 ml of 4% glucose in water and 0.5 ml of lysine free base, 200 mm. A stream of nitrogen gas at 7 L/min was flushed through the

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flask to evaporate chloroform over 5 minutes at 37 degrees

### Example 4

Synthesis of Reterovesicular Linconnes
Containing Granulocyte-Macrophase
Colony Stimulating Factor (GM-GSF)
Exactly the same procedure was used as in Example
2 except LP-2 was replaced with 1 up of GM-GSF.

## Example 5

Synthesis of Heterovesicular Liposomes of Various Lipid Composition, and Incorporation of Various Materials into Liposomes

In place of using diolecyl lecithin, dipalmtoyl

phosphatidylglyerol, cholesterol, and triolein (TO), and other amphipathic lipids such as phosphatidyl cholines (Pc). cardiolipin (CL), dimvristovl phosphatidvlglycerol (DMPG). phosphatidyl ethanolamines (PE), phosphatidyl serines (PS). dimyristoyl phosphatidic acid (DMPA), and other neutral lipids such as tricaprvlin (TC) in various combination can be used with similar results. For example, PC/C/CL/TO in 4.5/4.5/1/1 molar ration; DOPC/C/PS/TO in 4.5/4.5/1/1 molar ratio; PC/C/DPPG/TC in 5/4/1/1 molar ratio; PC/C/PG/TC in 5/4/1/1 molar ratio; PE/C/CL/TO in 4.5/4.5/1/1 molar ratio; and PC/C/DMPA/TO in 4.5/4.5/1/1 molar ratio can all be used. To incorporate other water-soluble materials, such as glucose, sucrose, methotrexate, Ponceau S, simply substitute the desired materials for IL-2 in Example 2. Also, other biologically active substances, such as set forth in Table 1, in suitable doses can be similarly substituted for II-2 as in Example 2.

### Example 6

In this example, the triolein in lipid components of above examples are substituted either singly or in combination by other triglycerides, vegetable oils, animal

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fats, tocopherols, tocopherol esters, cholesteryl esthers, or hydrocarbons with good results.

## Example 7

To make liposomes smaller than that in the foregoing examples, and with reference to Examples 1 or 2, the mechanical strength or duration of shaking or homogenization in Step 4 of Example 1 or 2 was increased. To make liposomes larger, the mechanical strength or duration of shaking or homogenization in Step 4 of Example 1 or 2 was decreased.

The heterovesicular liposomes can be administered to the patients in the normal manner when it is desirable to provide two separate biologically active compounds to the patient for the particular purpose of treatment desired.

The dosage range appropriate for human use includes the range of 1-6000 mg/s to body surface area. The reason that this range is so large is that for some applications, such as subcutaneous administration, the dose applications, such as subcutaneous administration, the dose desired to be used may be absolutely enormous. While doses outside the foregoing dose range may be given, this range encompasses the breadth of use for practically all the biologically active substances.

The multivesfcular liposomes may be administered by any desired route; for example, intrathecal, intraperitomeal, subcutamous, intravenous, intralymphatic, oral and submucomal, under many different kinds of epithelia including the bronchialar epithelia, the gastrointestinal epithelia, the urogenital epithelia, and various mucous membranes of the body, and intramsucular.

When encapsulating more than two substances separately in chambers of a liposome, a third (or fourth) aqueous component containing the third or fourth blologically active substance is formed, mixed to form a third or fourth water in oil smulston, and then combined

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with the first and second emulsions and mixed to form a "chimeric" emulsion containing the three or more biologically active substances. The remainder of the process is the same as described when encapsulating two biologically active compounds or substances.

The present invention, therefore, obtains the objects and ends and has the advantages mentioned as well as others inherent therein.

While examples of the invention have been given 10 for the purpose of disclosure, changes can be made therein which are within the spirit of the invention as defined by the appended claims.

What is claimed is:

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### Claims

- A heterovesicular lipid vesicle or liposome having at least two different substances, at least one of which is biologically active, encapsulated in separate chambers of the same liposome.
  - A process for producing a heterovesicular lipid vesicle or liposome having at least two different substances, at least one of which is biologically active, separately encapsulated in aqueous chambers thereof comprising the steps of:
    - (a) providing a first lipid component dissolved in one or more organic solvents and adding into the said lipid component an immiscible first aqueous component containing a first biologically active substance to be encapsulated;
    - (b) forming a first water-in-oil smulsion from the first two immiscible components; (c) providing a second lipid component dissolved in one or more organic solvents and adding into the said lipid component an immiscible second aqueous component containing a second
    - second aqueous component containing a second substance to be encapsulated; (d) forming a second water-in-cil emulsion from the second two immiscible components; (e) forming a chimeric emulsion by combining
    - the first water-in-oil emulsion and the second water-in-oil emulsion; (f) transferring and immersing the product of step (e) in a third media that is immiscible with said organic solvents;
    - (g) dispersing the chimeric emulsion to form solvent spherules containing multiple droplets of the first aqueous component containing the first substance and the second aqueous component containing the second substance; and

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(h) evaporating the organic solvents from the solvent spherules to form the heterovesicular liposomes.

- The process according to Claim 2 wherein the first and second lipid components are a phospholipid or an admixture of several phospholipids.
  - 4. The process according to Claim 2 wherein three or more water-in-oil emulsions containing three or more immiscible aqueous components are combined to form the chimeric emulsion.
  - The process according to Claim 2 wherein the first and second lipid components are identical.

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- 6. The process according to Claim 3 wherein the phospholipids are selected from the group consisting of phosphatidylcholine, cardiolipin, phosphatidylchanolamine, sphingomyelin, lysophosphatidylcholine, phosphatidylserine, phosphatidylinositol, phosphatidylgucerol, and phosphatidic acid.
- 7. The process according to Claim 3 wherein the one or more of the lipid components contain a lipid with a net negative charge or charges.
  - 8. The process according to Claim 3 wherein at least one of the phospholipids is provided in admixture with cholesterol.
- The process according to Claim 3 wherein at least one of the phospholipids is provided in admixture with stearylamine.

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- 10. The process according to Claim 2 wherein at least one of the first and second substances is a lipophilic biologically active material.
- 11. The process according to Claim 2 wherein at least one of the first and second lipid components is a neutral lipid either singly or in combination with a substance selected from the group consisting of triglycerides, vegetable oils, animal fats, tocopherols, tocopherol esterm, cholesteryl esterm, and hydrocarbons.
  - 12. The process according to Claim 2 wherein the organic solvent is selected from the group consisting of ethers, hydrocarbons, halogenated hydrocarbons, halogenated ethers, esters, and combinations thereof.
- 13. The process according to Claim 2 wherein the hydrochloride is selected from the group consisting of hydrochloric acid, lysine hydrochloride, histidine hydrochloride and combinations thereof.
  - 14. The process according to Claim 2 wherein the biologically active substance is hydrophilic.
  - 15. The process according to Claim 14 wherein the hydrophilic biologically active substance is selected from the group consisting of interleukin-2, cytosine arabinoside, methotrewate, 5-fluorouracil, cisplatin, floouridine, melphalan, merceptopurine, thioquanine; thioteps, vincristine, vinblastine, streptosocin, leuprolide, interferon, calcitonin, doxorubicin, daunorubicin, mitoxanthrone, amacrine, actinosycin, and bleomycin.
  - 16. The process according to Claim 2 wherein the emulsification of the two components is carried out using

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methods selected from the group consisting of mechanical agitation, ultrasonic energy, and nozzle atomization.

- 17. The process according to Claim 2 wherein the third aqueous component contains at least one acidneutralizing agent.
- 18. The process according to Claim 17 wherein the acid-neutralizing agent is selected either singly or in combination from the group consisting of free-base lysine and free-base histidine.
- The process according to Claim 2 wherein the third aqueous component has an ionic strength less than approximately 0.05.
- 20. The process according to Claim 17 wherein the third aqueous component is an aqueous solution further containing solutes selected from the group consisting of carbohydrates and aminoacids.
  - 21. The process according to Claim 17 wherein the third aqueous component is an aqueous solution containing solutes selected either singly or in combination from the group consisting of glucose, sucrose, lactose, free-base lysine, and free-base histidine.
    - 22. The process according to Claim 2 wherein the dispersion to form solvent spherules is carried out using methods selected from the group consisting of mechanical agitation, ultrasonic energy, and nozzle atomization.
    - 23. The process according to Claim 2 wherein the evaporation of the organic solvent is provided by passing nitrogen gas over the second aqueous component.

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- 24. The process of Claim 2 where, the biologically active substance to be encapsulated is selected from the group consisting of the compositions of Table 1.
- Heterovesicular liposomes made according to the method of Claim 2.
  - 26. A heterowesicular liposome containing at least two different substances, at least one of which is biologically active, encapsulated in separate chambers of the same liposome, at least one of the substances encapsulated in the presence of a hydrochloride.
  - 27. The process according to Claim 26 wherein the hydrochloride is selected from the group consisting of hydrochloric acid, lysine hydrochloride, histidine hydrochloride and combinations thereof.
- 28. A heterovesicalar liposome containing at least two substances of different compositions, at least one of which is biologically active, encapsulated in separate chambers of the liposome, at least one of the substances encapsulated in the presence of hydrochloric acid or other acid hydrochlorides and a neutralizing agent.
- 29. The heterovesicular liposome of Claim 26 where, the biologically active composition is selected from the group consisting of the compositions of Table 1.
- 30. The heterovesicular liposome of Claim 28 where, the biologically active composition is selected from the group consisting of the compositions of Table 1.
  - 31. A method for the treatment of a patient with two different substances, at least one of which is biologically active, comprising administering said

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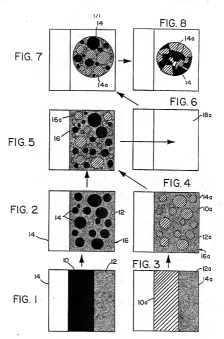
substances to the patient each encapsulated in separate chambers of a heterovesicular liposome.

- 32. A method for the treatment of a patient with at least two different substances, at least one of which is a biologically active compound, comprising administering said substances to the patient heterovesicular liposomes encapsulating the substances according to Claims 26, 27, 28, 29, 30 or 31.
- 33. A heterovesicular lipid vesicle or liposome 10 having at least two different substances, at least one of which is biologically active, encapsulated in separate chambers of the same liposome where,

the biologically active substances are selected from the group consisting of antiarrhythmic, antiasthms, antibiotic, anticancer, antialabetic, antifungal, antihistanines, antihypertensives, antihypotension, antiparasitic, antiviral, cell surface receptor blockers, cardiac glycosides, hormones, immunoptherapies, nucleic acids and analogs, proteins and glycoproteins, sedatives and analogs, proteins and glycoproteins, sedatives and analgesic, steroids, tranquilizers, and vaccines.

34. The process of claim 2 where.

the biologically active substances are selected from the group consisting of antiarrhythmic, antiasthma, antibiotic, anticancer, antidiabetic, antifungal, antihistamines, antihypertensives, antihiptotension, antiparasitic, antiviral, cell surface receptor blockers, cardiac glycosides, hormones, immunoptherapies, nucleic acids and analogs, proteins and glycoproteins, sedatives and analogs, proteins and glycoproteins, and vecoines.



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